# Prediction of Medical Expenses of Individuals using Regression Models

### Parvez Khan

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### **Dataset**

Here we have a dataset about medical costs billed by health insurance on different individuals along with their age, sex, bmi, number of children and other parameters collected from kaggle database.

### **Objective**

To predict the medical bill for individuals based on different parameters using different regression models and choose the best among them.

### **Data description**

Loading the data

```
data <- read.csv("insurance.csv", header = TRUE)</pre>
```

Checking for number of rows and columns

```
dim(data)
## [1] 1338 7
```

Taking a look at the data frame

```
head(data)
##
                  bmi children smoker
    age
           sex
                                       region
                                                charges
## 1 19 female 27.900
                                yes southwest 16884.924
## 2 18
          male 33.770
                            1
                                 no southeast 1725.552
          male 33.000
                                 no southeast 4449.462
## 3 28
                            3
## 4 33
          male 22.705
                           0
                                 no northwest 21984.471
          male 28.880
                           0
                           0
## 5 32
                                 no northwest
                                               3866.855
## 6 31 female 25.740
                                 no southeast
                                               3756,622
str(data)
## 'data.frame':
                  1338 obs. of 7 variables:
   $ age : int 19 18 28 33 32 31 46 37 37 60 ...
## $ sex
             : chr "female" "male" "male" ...
## $ bmi
             : num 27.9 33.8 33 22.7 28.9 ...
## $ children: int 0 1 3 0 0 0 1 3 2 0 ...
## $ smoker : chr "yes" "no" "no" "no" ...
## $ region : chr "southwest" "southeast" "southeast" "northwest" ...
## $ charges : num 16885 1726 4449 21984 3867 ...
```

### **Data Cleaning**

Checking for missing values

```
missing <- sum(is.na(data))
missing
## [1] 0</pre>
```

**Comment:** There is no missing value in the data.

Checking for duplicate values

```
duplicate_rows <- data[duplicated(data),]
duplicate_rows

## age sex bmi children smoker region charges
## 582 19 male 30.59 0 no northwest 1639.563</pre>
```

**Comment:** There is one duplicate row, so we remove it from the data.

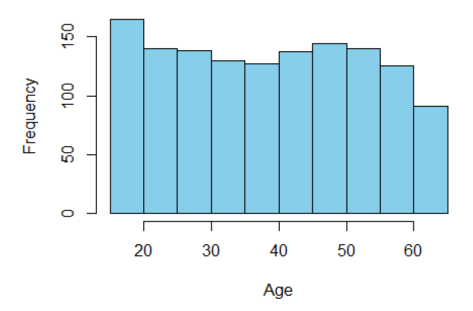
```
data <- data[!duplicated(data),]</pre>
```

#### **Data Visualization**

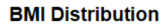
To start with data visualization we first plot histograms for the numeric columns

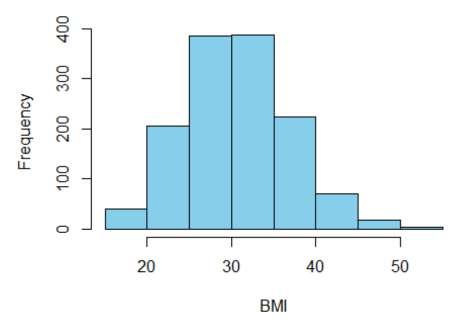
```
hist(data$age, main = "Age Distribution", xlab = "Age", ylab = "Frequency",
col = "skyblue")
```

# Age Distribution



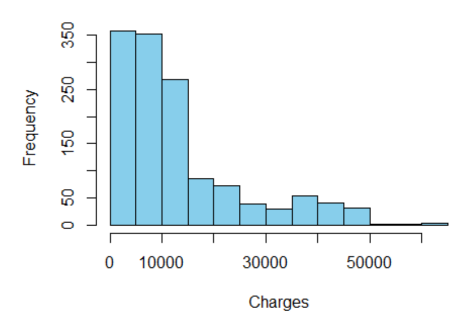
hist(data\$bmi, main = "BMI Distribution", xlab = "BMI", ylab = "Frequency",
col = "skyblue")





```
hist(data$charges, main = "Charges Distribution", xlab = "Charges", ylab =
"Frequency", col = "skyblue")
```

### **Charges Distribution**



Now for the categorical columns first we will get the number of occurrence for each unique entries

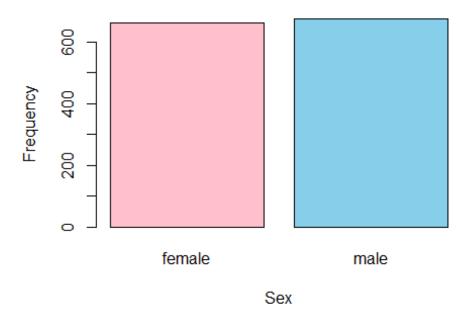
```
sex_count <- table(data$sex)</pre>
sex_count
##
## female
             male
      662
              675
##
children_count <- table(data$children)</pre>
children_count
##
##
     0
         1
              2
                  3
                      4
                           5
## 573 324 240 157 25 18
smoker_count <- table(data$smoker)</pre>
smoker_count
##
##
     no yes
## 1063 274
```

```
region_count <- table(data$region)
region_count
##
## northeast northwest southeast southwest
## 324 324 364 325</pre>
```

Now to visualize the distribution of categorical columns we will plot their bar graph

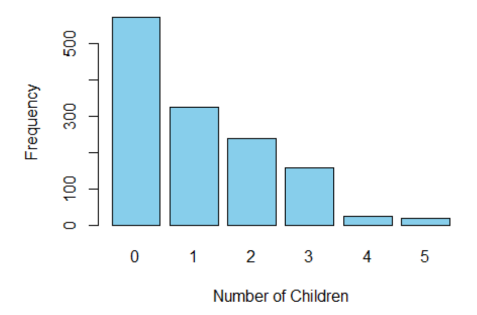
```
barplot(sex_count, main = "Sex Distribution", xlab = "Sex", ylab =
"Frequency", col = c("pink", "skyblue"))
```

### **Sex Distribution**



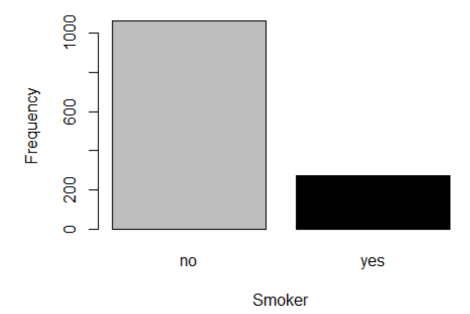
```
barplot(children_count, main = "Children Distribution", xlab = "Number of
Children", ylab = "Frequency", col = "skyblue")
```

### **Children Distribution**



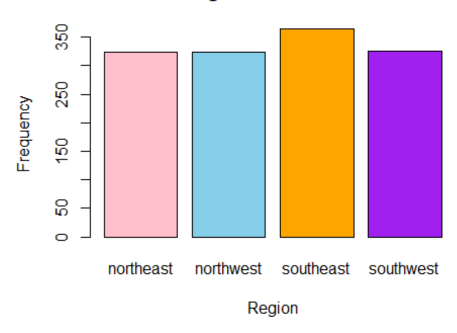
barplot(smoker\_count, main = "Smoker Distribution", xlab = "Smoker", ylab =
"Frequency", col = c("grey", "black"))

### **Smoker Distribution**



```
barplot(region_count, main = "Region Distribution", xlab = "Region", ylab =
"Frequency", col = c("pink", "skyblue", "orange", "purple"))
```

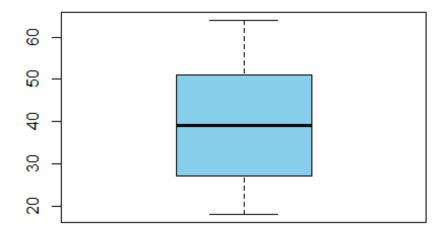
## **Region Distribution**



Now we plot the boxplot for numerical columns to check for outliers

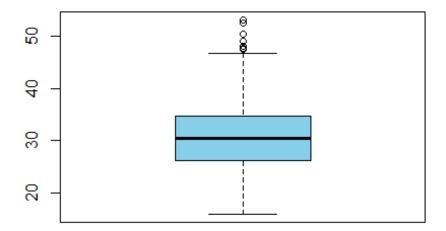
```
boxplot(data$age, main = "Boxplot for Age", col = "skyblue")
```

# **Boxplot for Age**



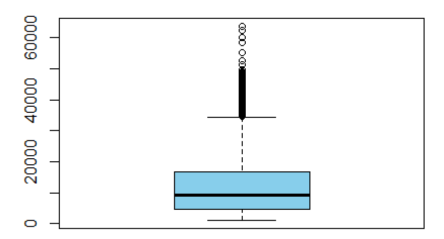
boxplot(data\$bmi, main = "Boxplot for BMI", col = "skyblue")

# **Boxplot for BMI**



boxplot(data\$charges, main = "Boxplot for Charges", col = "skyblue")

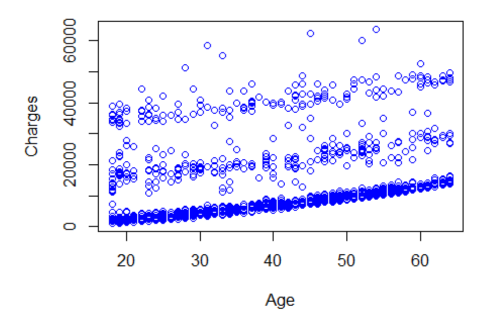
## **Boxplot for Charges**



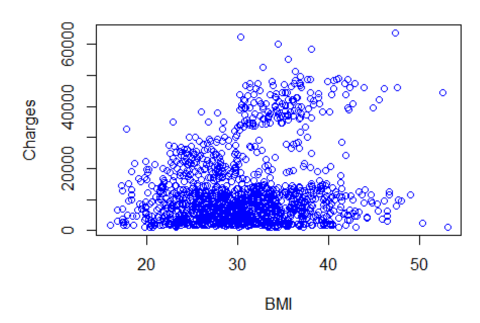
**Comment:** It is evident from the boxplot that bmi and chrages contains some outliers while there is no outliers on the age column.

Now we plot a scatter diagram for further interpretation

```
plot(data$age, data$charges, xlab = "Age", ylab = "Charges", col = "blue")
```



plot(data\$bmi, data\$charges, xlab = "BMI", ylab = "Charges", col = "blue")



**Comment:** From the scatter plot it can be seen that their is a weak relation between age and charges while the relation is more weak between bmi and charges.

Now we create a correlation matrix to further check for quantified relationship between the variables.

```
numeric_data <- data[, c("age", "bmi", "charges")]
cor(numeric_data)

## age bmi charges

## age 1.0000000 0.1093436 0.2983082

## bmi 0.1093436 1.0000000 0.1984008

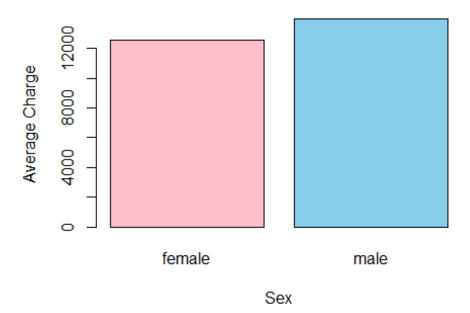
## charges 0.2983082 0.1984008 1.0000000
```

**Comment:** From the correlation matrix it is evident that the correlation between bmi and charges is very low while between age and charges it is slightly better.

Now we see if the charges vary with different categories

Charges for males and females:

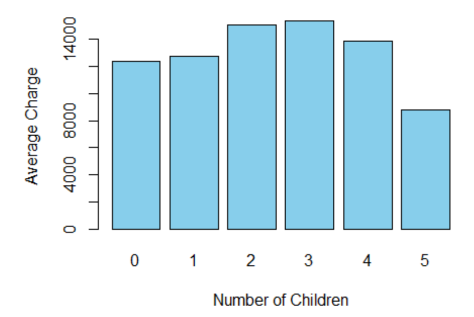
```
av_sex_charge <- tapply(data$charges, data$sex, mean)
barplot(av_sex_charge, xlab = "Sex", ylab = "Average Charge", col = c("pink",
"skyblue"))</pre>
```



**Comment:** As it can be seen there is not much difference in medical charge of male and female.

Charges for different number children:

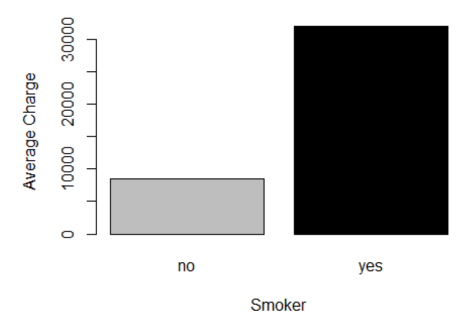
```
av_children_charge <- tapply(data$charges, data$children, mean)
barplot(av_children_charge, xlab = "Number of Children", ylab = "Average
Charge", col = "skyblue")</pre>
```



**Comment:** From the plot we can see that the average charge is almost not very different among people with different number of children with an exception where surprisingly people with 5 children has less average medical charge.

Charges for smokers and non-smokers:

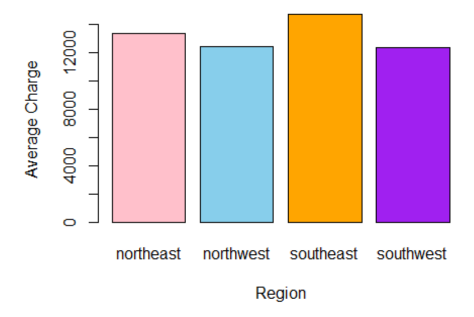
```
av_smoker_charge <- tapply(data$charges, data$smoker, mean)
barplot(av_smoker_charge, xlab = "Smoker", ylab = "Average Charge", col =
c("grey", "black"))</pre>
```



**Comment:** It is clearly evident from the plot that people who smoke tends to have a significantly higher average medical cost than people who don't.

Charges for poeple of different regions:

```
av_region_charge <- tapply(data$charges, data$region, mean)
barplot(av_region_charge, xlab = "Region", ylab = "Average Charge", col =
c("pink", "skyblue", "orange", "purple"))</pre>
```



**Comment:** Here also we see that there is not much difference in charges among people from different region.

### **Model Fitting**

Before we fit any model into the data, at first we need to change the categorical columns into numerical ones so that we can work with them with ease.

```
data$sex <- as.numeric(factor(data$sex)) - 1</pre>
data$smoker <- as.numeric(factor(data$smoker)) - 1</pre>
data$region <- as.numeric(factor(data$region)) - 1</pre>
head(data)
##
                 bmi children smoker region
                                               charges
     age sex
                                           3 16884.924
## 1
     19
           0 27.900
                            0
                                    1
           1 33.770
## 2
      18
                            1
                                    0
                                           2 1725.552
## 3 28
           1 33.000
                            3
                                    0
                                           2 4449.462
           1 22.705
                            0
     33
                                    0
                                           1 21984.471
## 4
## 5
     32
           1 28.880
                            0
                                    0
                                           1 3866.855
           0 25.740
## 6 31
                            0
                                    0
                                           2 3756.622
```

We have now changed the whole dataset into numerical data and can now proceed with model fitting.

### Split the data into training and testing

Here we allocate 80% of the data for training and the remaining 20% goes for testing.

```
set.seed(82)
indices <- sample(nrow(data), 0.8*nrow(data))</pre>
train_data <- data[indices, ]</pre>
test_data <- data[-indices, ]</pre>
```

*Linear Regression Model:* 

```
lr_model <- lm(charges ~ ., data = train_data)</pre>
summary(lr_model)
##
## Call:
## lm(formula = charges ~ ., data = train data)
## Residuals:
##
     Min
            1Q Median
                          3Q
                               Max
## -11288 -2916 -1061
                        1382 29786
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
13.52 19.336 < 2e-16 ***
                261.46
## age
## sex
                 16.74
                          379.25 0.044 0.96479
                           31.27 10.557 < 2e-16 ***
## bmi
                330.14
## children
               453.81
                           153.75 2.952 0.00323 **
              23539.33 464.88 50.635 < 2e-16 ***
-344.47 170.67 -2.018 0.04381 *
## smoker
## region
              -344.47
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6169 on 1062 degrees of freedom
## Multiple R-squared: 0.7439, Adjusted R-squared: 0.7425
## F-statistic: 514.3 on 6 and 1062 DF, p-value: < 2.2e-16
```

**Comment:** From the summary we can see that the p-value for the coefficient of the variable 'sex' is 0.96 which is significantly high(>>0.05), specifying that it doesn't affect the response variable 'charges'. So now we will rebuild the model dropping the variable 'sex'.

```
lr model2 <- lm(charges ~ . -sex, data = train data)</pre>
summary(lr model2)
##
## Call:
## lm(formula = charges ~ . - sex, data = train_data)
##
## Residuals:
##
     Min
            10 Median
                         3Q
                              Max
## -11280 -2914 -1068
                       1391 29793
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
13.50 19.363 < 2e-16 ***
                261.44
## age
                          31.24 10.569 < 2e-16 ***
## bmi
                330.18
                         153.67 2.953 0.00321 **
## children
              453.87
                        463.03 50.841 < 2e-16 ***
              23541.05
## smoker
## region
              -344.54
                         170.59 -2.020 0.04366 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 6166 on 1063 degrees of freedom
## Multiple R-squared: 0.7439, Adjusted R-squared: 0.7427
## F-statistic: 617.7 on 5 and 1063 DF, p-value: < 2.2e-16
```

#### Model Accuracy Check:

Now we will check the accuracy of the model and check how it performs with the test data

```
lr_train_predicted <- predict(lr_model2, newdata = train_data)
lr_test_predicted <- predict(lr_model2, newdata = test_data)

rmse <- function(actual, predicted){
    sqrt(mean((actual - predicted)^2))
}

rsquared <- function(actual, predicted){
    sst <- sum((actual - mean(actual))^2)
    ssr <- sum((actual - predicted)^2)
    1 - (ssr/sst)
}

lr_rmse_train <- rmse(train_data$charges, lr_train_predicted)
lr_rsquared_train <- rsquared(train_data$charges, lr_train_predicted)

lr_rmse_test <- rmse(test_data$charges, lr_test_predicted)
lr_rsquared_test <- rsquared(test_data$charges, lr_test_predicted)

cat("Trainig RMSE: ", lr_rmse_train, "\nTraining R-squared: ",</pre>
```

```
lr_rsquared_train, "\n\nTesting RMSE: ", lr_rmse_test, "\nTesting R-squared:
", lr_rsquared_test, "\n")
## Trainig RMSE: 6148.358
## Training R-squared: 0.7439435
##
## Testing RMSE: 5629.571
## Testing R-squared: 0.7754724
```

### **Decision Tree Regression Model:**

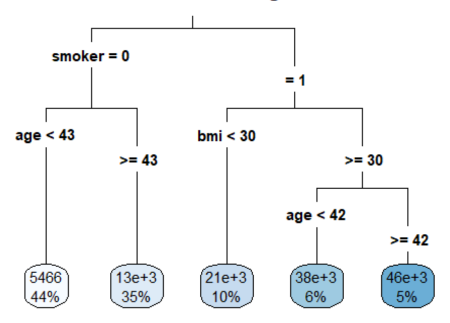
First we load the library 'rpart' to perform Decision Tree regression

```
library(rpart)
library(rpart.plot)
```

Now we train our decision tree model with the train data and visualize it

```
dt_model <- rpart(charges ~ ., data = train_data, method = "anova")
rpart.plot(dt_model, type = 3, main = "Decision Tree for Charge Prediction")</pre>
```

### **Decision Tree for Charge Prediction**



### Model Accuracy Check:

Now we check the performance of the model with the test data

```
dt_train_predicted <- predict(dt_model, newdata = train_data)
dt_test_predicted <- predict(dt_model, newdata = test_data)</pre>
```

```
dt_rmse_train <- rmse(train_data$charges, dt_train_predicted)
dt_rsquared_train <- rsquared(train_data$charges, dt_train_predicted)

dt_rmse_test <- rmse(test_data$charges, dt_test_predicted)
dt_rsquared_test <- rsquared(test_data$charges, dt_test_predicted)

cat("Training RMSE: ", dt_rmse_train, "\nTraining R-squared: ",
dt_rsquared_train, "\n\nTesting RMSE: ", dt_rmse_test, "\nTesting R-squared:
", dt_rsquared_test, "\n")

## Training RMSE: 4988.915
## Training R-squared: 0.8314108
##
## Testing RMSE: 4472.26
## Testing R-squared: 0.8582989</pre>
```

#### Random Forest Regression Model:

First we load the library 'randomForest' to perform random forest regression

```
library(randomForest)
## randomForest 4.7-1.1
## Type rfNews() to see new features/changes/bug fixes.
```

Now we train the random forest model with the train data

```
rf_model <- randomForest(charges ~ ., data = train_data)
print(rf_model)

##
## Call:
## randomForest(formula = charges ~ ., data = train_data)
## Type of random forest: regression
## Number of trees: 500
## No. of variables tried at each split: 2
##
## Mean of squared residuals: 23814785
## War explained: 83.87</pre>
```

### Model Accuracy Check

```
rf_train_predicted <- predict(rf_model, newdata = train_data)
rf_test_predicted <- predict(rf_model, newdata = test_data)

rf_rmse_train <- rmse(train_data$charges, rf_train_predicted)
rf_rsquared_train <- rsquared(train_data$charges, rf_train_predicted)

rf_rmse_test <- rmse(test_data$charges, rf_test_predicted)
rf_rsquared_test <- rsquared(test_data$charges, rf_test_predicted)</pre>
```

```
cat("Training RMSE: ", rf_rmse_train, "\nTraining R-squared: ",
rf_rsquared_train, "\n\nTesting RMSE: ", rf_rmse_test, "\nTesting R-squared:
", rf_rsquared_test, "\n")
## Training RMSE: 3126.354
## Training R-squared: 0.9337945
##
## Testing RMSE: 4191.63
## Testing R-squared: 0.8755242
```

#### **Conclusion**

After fitting and checking the performance of all the models, We can see that the accuracy in training is 74%, 83%, and 93% for linear regression, decision tree and random forest model respectively, specifying the superiority of the random forest model over the other two models while training with the train data. On the other hand in terms of predicting, the accuracy of the linear regression model and the decision tree model slightly increases to 77% and 86% respectively, while the accuracy of the random forest model decreases to 87% which is still better than the other two models. So we come to the conclusion that **Random Forest Model** is the best among all the models.